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APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.	
10/638,173	08/06/2003	Robert Kain	ILLINC.026C1	3813	
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Please find below and/or attached an Office communication concerning this application or proceeding.

The time period for reply, if any, is set in the attached communication.

Notice of the Office communication was sent electronically on above-indicated "Notification Date" to the following e-mail address(es):

jcartee@kmob.com eOAPilot@kmob.com

Application No. Applicant(s) 10/638,173 KAIN ET AL. Office Action Summary Examiner Art Unit BJ Forman 1634 -- The MAILING DATE of this communication appears on the cover sheet with the correspondence address --Period for Reply A SHORTENED STATUTORY PERIOD FOR REPLY IS SET TO EXPIRE 3 MONTH(S) OR THIRTY (30) DAYS. WHICHEVER IS LONGER, FROM THE MAILING DATE OF THIS COMMUNICATION. Extensions of time may be available under the provisions of 37 CFR 1.136(a). In no event, however, may a reply be timely filed after SIX (6) MONTHS from the mailing date of this communication. - If NO period for reply is specified above, the maximum statutory period will apply and will expire SIX (6) MONTHS from the mailing date of this communication. Failure to reply within the set or extended period for reply will, by statute, cause the application to become ABANDONED (35 U.S.C. § 133). Any reply received by the Office later than three months after the mailing date of this communication, even if timely filed, may reduce any earned patent term adjustment. See 37 CFR 1.704(b). Status 1) Responsive to communication(s) filed on 17 December 2008. 2a) ☐ This action is FINAL. 2b) This action is non-final. 3) Since this application is in condition for allowance except for formal matters, prosecution as to the merits is closed in accordance with the practice under Ex parte Quayle, 1935 C.D. 11, 453 O.G. 213. Disposition of Claims 4) Claim(s) 60.65-71.76-83.88-96.101-117 and 126-133 is/are pending in the application. 4a) Of the above claim(s) _____ is/are withdrawn from consideration. 5) Claim(s) _____ is/are allowed. 6) Claim(s) 60,65-71,76-83,88-96,101-117 and 126-133 is/are rejected. 7) Claim(s) _____ is/are objected to. 8) Claim(s) _____ are subject to restriction and/or election requirement. Application Papers 9) The specification is objected to by the Examiner. 10) ☐ The drawing(s) filed on _____ is/are: a) ☐ accepted or b) ☐ objected to by the Examiner. Applicant may not request that any objection to the drawing(s) be held in abeyance. See 37 CFR 1.85(a). Replacement drawing sheet(s) including the correction is required if the drawing(s) is objected to. See 37 CFR 1.121(d). 11) The oath or declaration is objected to by the Examiner. Note the attached Office Action or form PTO-152. Priority under 35 U.S.C. § 119 12) Acknowledgment is made of a claim for foreign priority under 35 U.S.C. § 119(a)-(d) or (f). a) ☐ All b) ☐ Some * c) ☐ None of: Certified copies of the priority documents have been received. 2. Certified copies of the priority documents have been received in Application No. Copies of the certified copies of the priority documents have been received in this National Stage application from the International Bureau (PCT Rule 17.2(a)). * See the attached detailed Office action for a list of the certified copies not received. Attachment(s) 1) Notice of References Cited (PTO-892) 4) Interview Summary (PTO-413) Paper No(s)/Mail Date. Notice of Draftsperson's Patent Drawing Review (PTO-948)

Information Disclosure Statement(s) (PTO/SB/08)
 Paper No(s)/Mail Date 12/08.

5) Notice of Informal Patent Application

6) Other:

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DETAILED ACTION

Continued Examination Under 37 CFR 1.114

1. A request for continued examination under 37 CFR 1.114, including the fee set forth in 37 CFR 1.17(e), was filed in this application after final rejection. Since this application is eligible for continued examination under 37 CFR 1.114, and the fee set forth in 37 CFR 1.17(e) has been timely paid, the finality of the previous Office action has been withdrawn pursuant to 37 CFR 1.114. Applicant's submission filed on 17 December 2008 has been entered.

Status of the Claims

This action is in response to papers filed 17 December 2008 in which claims 60,
 65, 66, 71, 76, 77, 83, 88, 89 and 94 were amended, claims 61, 63, 72, 74, 84, 86, 97,
 99 and 118-125 were canceled and claims 127-133 were added. The amendments have been thoroughly reviewed and entered.

The previous rejections in the Office Action dated 26 June 2008 are withdrawn in view of the amendments. Applicant's arguments have been thoroughly reviewed but are deemed moot in view of the amendments, withdrawn rejections and new grounds for rejection. New grounds for rejection are discussed.

Claims 60, 65-71, 76-83, 88-96, 101-117 and 126-133 are under prosecution.

Claim Rejections - 35 USC § 103

- 3. The following is a quotation of 35 U.S.C. 103(a) which forms the basis for all obviousness rejections set forth in this Office action:
 - (a) A patent may not be obtained though the invention is not identically disclosed or described as set forth in section 102 of this title, if the differences between the

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subject matter sought to be patented and the prior art are such that the subject matter as a whole would have been obvious at the time the invention was made to a person having ordinary skill in the art to which said subject matter pertains. Patentability shall not be negatived by the manner in which the invention was made.

Claims 60, 66-71, 77-83, 89-96, 103-104, 106-117, 127, 129 and 131-132 are rejected under 35 U.S.C. 103(a) as being unpatentable over Walt et al. (U.S. Patent No. 6,327,410, filed 11 September 1998) in view of McDevitt et al. (U.S. Patent No. 6,680,206, filed 16 July 1999) and/or Seul et al. (U.S. Patent No. 7,041,510, filed 17 September 1999).

Regarding Claim 60, 71, 83, 94, Walt et al. disclose an array and method of making the array comprising a substrate having a surface (Column 5, lines 32-60), a first assay location and second assay location on the surface (Column 5, line 61-Column 6, line 30), wherein the substrate has a first plurality of depressions in first and second assay locations and first and second microsphere populations randomly placed in the assay locations wherein the assay locations spatially identifiable (Column 18, line 59-Column 18, line 5).

Walt et al teach the assay locations comprising marker beads (Column 19, line 4) but do not teach blank beads in the assay locations.

However, blank beads were well known and routinely practiced in the art of bead arrays at the time the invention was made as taught by McDevitt and Seul.

McDevitt et al teach a similar array and method of making the array comprising a substrate having a surface (Fig. 3), a first assay location and second assay location on the surface (#250, Fig. 3, Column 39, lines 15-34 and Column 40, lines 34-37), a first

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plurality of depressions in first and second assay locations and first and second microsphere populations randomly placed in the assay locations (Column 10, lines 13-16) and wherein assay locations comprises blank microspheres (Column 25, lines 1-45 and Fig. 16) wherein the blank microspheres provide a reference signal to which multiple different signals can be compared thereby allowing simultaneous evaluation of multiple chemically distinct analytes (Column 25, lines 1-8).

Seul also teach a similar array and method of making the array comprising a substrate having multiple assay locations and multiple populations of beads randomly distributed in the assay locations (e.g. Fig. 28) wherein the assay locations further comprise blank beads (i.e. spacer particles) to provide interparticle spacing of analyte beads and thereby allowing optical analysis of individual analyte particles (Column 25, lines 2-21).

It would have been obvious to one of ordinary skill in the art at the time the claimed invention was made to apply the blank beads of McDevitt and/or Seul to the array of Walt. One of ordinary skill in the art would have been motivated to do so with a reasonable expectation of success and for the added benefits of providing appropriate spacing of analyte beads allowing optical analysis of individual analyte particles (Seul, Column 25, lines 2-21) and/or providing a reference signal to which multiple different signals can be compared allowing simultaneous evaluation of multiple chemically distinct analytes (McDevitt, Column 25, lines 1-8).

Regarding Claim 66, 77, 89, Walt et all teach the array wherein the bioactive agent comprises a nucleic acid (Column 10, lines 28-35). McDevitt et all disclose the

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array wherein the bioactive agent comprises a nucleic acid (Column 5, lines 45-65) and Seul et al. teach the bioactive agent comprises a nucleic acid (Example IX).

Regarding Claims 68, 79, 91, 103, Walt et al. teach the array is within a hybridization chamber (Fig. 4). McDevitt et al. teach the similar array wherein the substrate is enclosed within a hybridization chamber (Fig. 17, Column 26-27).

Regarding Claims 69, 80, 92, 104, Walt et al teach the array wherein the substrate comprises a membrane i.e. over the beads (Column 6, lines 45-47) and McDevitt et al teach the similar array wherein the hybridization chamber comprises a flexible membrane (Column 11, line s 40-44).

Regarding Claims 82 and 106, Walt et al teach the depressions are wells (Column 6, lines 16-30). McDevitt et al teach the similar array wherein the depressions are wells (Fig. 3) and Seul et al teach the similar array wherein the depressions are wells formed via hydrophobic grid (Column 23, lines 9-15).

Regarding Claim 107-108, Walt et al teach the method further comprising preparation of the DNA by PCR (Column 23, lines 5-8).

Regarding Claim 109-111, Walt et al. teach the method wherein the bioactive agent is DNA (Column 10, lines 28-35) and the method includes sequencing (Column 24, lines 51-52). While Walt et al. teaches the array is used for sequencing, the sequencing practiced with the array produced by the method, does not further define the method of making the array. As such, the recited sequencing methods do not further define the method of Claim 94 for making the array. Furthermore, Felder et al teaches the similar method determines the sequence (column 11, lines 23-48).

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Regarding Claims 95-96, 112-117, Walt et al. teach the arrays and methods wherein each subpopulation is randomly distributed such that members of each subpopulation are in multiple sub-bundles (Column 18, line 48-Column 19, line 53). Seul et al. teach the similar array wherein each subpopulation is randomly distributed such that members of each subpopulation are in each assay location (Fig. 28, Column 44, lines 43-67).

Regarding Claims 126, 129, 131 and 132, Seul et al teach the similar array wherein hybridization between the target and bean-immobilized capture probe occur prior to bead immobilization on the support thereby providing both double and single stranded nucleic acids on the beads. Seul teaches that hybridization prior to surface immobilization facilitate subsequent analysis of strands of interest (Column 32, line 62-Column 33, line 7). It would have been obvious to one of ordinary skill in the art at the time the claimed invention was made to apply the solution hybridization prior to immobilization as taught by Seul to the array and method of Walt. One of ordinary skill in the art would have been motivated to do so with a reasonable expectation of success and for the added benefit of facilitating subsequent analysis of strands of interest as desired in the art (Seul, Column 32, line 62-Column 33, line 7).

Claims 65, 67, 70, 76, 78, 81, 88, 90, 93, 101, 102, 105, 126, 128, 130, 133 are rejected under 35 U.S.C. 103(a) as being unpatentable over Walt et al. (U.S. Patent No.

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6,327,410, filed 11 September 1998) in view of McDevitt et al. (U.S. Patent No. 6,680,206, filed 16 July 1999) and/or Seul et al. (U.S. Patent No. 7,041,510, filed 17 September 1999) as applied to Claims 60, 71, 83, 94 above and further in view of Felder et al. (U.S. Patent No. 6,232,066, filed 2 July 1998).

Regarding Claims 65, 70, 76, 81, 88, 93, 101, 105, 126, 130, 133, Walt et al teach the assay locations are spatially identifiable manually but the reference does not specifically teach the assay locations are separated. However, array locations separated by gaskets were well known in the art at the time the claimed invention was made as taught by Felder et al (Fig. 4-5).

Felder et al teach a substrate (Column 5, lines 1-13) having a plurality of assay locations (regions), each having a subpopulation of bioactive agents (e.g. genomic DNA, Column 6, lines 52-67) wherein the assay locations are separated by a gasket forming an array of wells-within-wells (e.g. wax or silicone barriers/well separator, Column 5, lines 19-59; Column 6, lines 38-51; Column 13, lines 1-22; and Fig. 5) whereby the assay locations are spatially discrete, identifiable and addressable within a fluidically controlled environment (Column 5, lines 19-59).

Walt clearly desires segregation of the subpopulations to provide spatial encoding of the microspheres and suggests manual techniques to do so (Column 18, line 59-Column 19, line 5). It would have been obvious to one of ordinary skill in the art at the time the claimed invention was made to apply the barrier elements of Felder et al to the substrate of Walt et al. One of ordinary skill in the art would have been motivated to do so based on the desired segregation of Walt et al. and further for the

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expected benefit of providing for fluidically controlled multi-sample testing without cross contamination between adjacent regions as taught by Felder et al (Column 5, lines 19-59).

Regarding Claim 67, 78, 90, 102, Walt et al. teach the array wherein the support is planar glass (Column 5, lines 57-60). McDevitt (Column 8, line 64) and Seul (Column 9, line 46) also teach glass substrate. Felder et al. teach the similar array wherein the glass support is a glass slide (Column 5, line 2). It would have been obvious to one of ordinary skill in the art at the time the claimed invention was made to apply the slide of Felder to the glass substrates of Walt, McDevitt and/or Seul. One of ordinary skill in the art would have been motivated to do so based on the commercial availability of microscope slides.

Conclusion

6. No claim is allowed.

Any inquiry concerning this communication or earlier communications from the examiner should be directed to BJ Forman whose telephone number is (571) 272-0741. The examiner can normally be reached on 6:00 TO 3:30.

If attempts to reach the examiner by telephone are unsuccessful, the examiner's supervisor, Ram Shukla can be reached on (571) 272-0735. The fax phone number for the organization where this application or proceeding is assigned is 571-273-8300.

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Information regarding the status of an application may be obtained from the Patent Application Information Retrieval (PAIR) system. Status information for published applications may be obtained from either Private PAIR or Public PAIR. Status information for unpublished applications is available through Private PAIR only. For more information about the PAIR system, see http://pair-direct.uspto.gov. Should you have questions on access to the Private PAIR system, contact the Electronic Business Center (EBC) at 866-217-9197 (toll-free). If you would like assistance from a USPTO Customer Service Representative or access to the automated information system, call 800-786-9199 (IN USA OR CANADA) or 571-272-1000.

BJ Forman Primary Examiner Art Unit 1634

/BJ Forman/ Primary Examiner, Art Unit 1634